## Amendments to the Claims:

 (Previously presented) An antigen composition capable of eliciting an enhanced cytotoxic T cell response in the context of a major histocompatibility complex class I molecule (MHC class I), comprising a soluble protein antigen modified by chemical linkage or fusion to an added peptidic sequence consisting of SEQ ID NO:4, wherein said added peptidic sequence facilitates entry of said antigen into antigen presenting cells (APC).

## 2-4. (Cancelled)

- (Previously presented) The antigen composition of claim 1 wherein said antigen is specific to a tumor or pathogen.
- (Previously presented) The antigen composition of claim 1, wherein said added peptidic sequences is covalently linked to said antigen.
- (Previously presented) The antigen composition of claim 1 wherein said antigen composition is a fusion protein produced by translation of a continuous nucleotide coding sequence.

## 8-18. (Cancelled)

19. (New) A therapeutic composition, comprising an antigen presenting cell (APC), stimulated by exposure in vitro to a modified antigen having an added peptidic sequence which

antigen into APC, wherein said stimulated APC is effective to activate T-cells to produce a cytotoxic cellular immune response against said antigen, at a T-cell activation level that is higher than that produced by such APC stimulated by the antigen alone.

- 20. (New) The therapeutic composition of claim 19, wherein said added peptidic sequence comprises one or more sequences selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5. SEQ ID NO:6 and SEQ ID NO:7.
- 21. (New) The therapeutic composition of claim 19, wherein said added peptidic sequence comprises a sequence presented as CYS-[X-Y-Y-Y-Y]n; wherein X= glu or asp, Y = ala, leu, ile, phe, gly, cys, met or val and n is greater than or equal to 3 or [X-Y-Y-Y-Y-Y]n; wherein X= glu or asp, Y = ala, leu, ile, phe, gly, cys, met or val and n is greater than or equal to 3.
- (New) The therapeutic composition of claim 19, wherein said antigen is a soluble protein antigen.
- 23. (New) The therapeutic composition of claim 19, for use in immunizing a subject against a tumor or pathogen wherein said antigen is specific to the tumor or pathogen.
- 24. (New) The therapeutic composition of claim 19, wherein said one or more added peptidic sequences are covalently linked to said antigen.
- 25. (New) The therapeutic composition of claim 19, wherein said antigen is a fusion protein produced by translation of a continuous nucleotide coding sequence.
- 26. (New) A method of immunizing a subject against a tumor or pathogen having a known cancer- or pathogen-specific antigen, comprising:
  - (a) obtaining a blood sample from a subject;
  - (b) isolating a monocyte fraction from said blood sample;
  - (c) enriching said monocyte fraction to obtain a population of DC;

- (d) pulsing said DC in vitro with a selected antigen composition comprising a selected antigen and an added peptidic sequence which facilitates entry of said antigen into antigen presenting cells (APC) by exposing said DC to the antigen composition in a manner effective to induce cell-surface presentation of one or more peptide antigens against which an immune response is desired; and
  - (e) returning the pulsed DC to the subject.
- 27. (New) The immunization method of claim 26, wherein said selected antigen is a cancer- or pathogen-specific antigen.
- 28. (New) The immunization method of claim 26, for use in treating cancer wherein said selected antigen is a cancer-specific antigen.
- (New) The immunization method of claim 28, further comprising administering to the patient an anti-cancer agent.
- 30. (New) The antigen composition of claim 1, wherein said added peptidic sequence comprises one or more sequences selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6 and SEQ ID NO:7.
- 31. (New) The antigen composition of claim 1, wherein said added peptidic sequence comprises a sequence presented as CYS-[X-Y-Y-Y-Y]n; wherein X= glu or asp, Y = ala, leu, ile, phe, gly, cys, met or val and n is greater than or equal to 3 or [X-Y-Y-Y-Y]n; wherein X= glu or asp, Y = ala, leu, ile, phe, gly, cys, met or val and n is greater than or equal to 3.